

### **PCT**

## WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau

#### INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification <sup>6</sup> :		(11) International Publication Number: WO 98/3	1550
B41M 5/36, B41C 1/10	A1	(43) International Publication Date: 23 July 1998 (23.	07.98)
(21) International Application Number: PCT/EP( (22) International Filing Date: 8 January 1998 (6)		DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NI	
(30) Priority Data: 97200144.0 17 January 1997 (17.01.97) (34) Countries for which the regional or international application was filed:	DE et	Before the expiration of the time limit for amendi	
(71) Applicant: AGFA-GEVAERT NAAMLOZE NOOTSCHAP [BE/BE]; Septestraat 27, B-2640 (BE).		•	
(72) Inventors: GRABLEY, Fritz-Feo; Agfa-Gevae PRO/IIE 3804, Septestraat 27, B-2640 Morts GRIES, Willi-Kurt; Agfa-Gevaert N.V., PRO/I Septestraat 27, B-2640 Mortsel (BE). SCHI Hans-Joachim; Agfa-Gevaert N.V., PRO/IIE 3804 traat 27, B-2640 Mortsel (BE).	iel (BI IE 380 LOSSE		
(54) Title: LASER-IMAGEABLE RECORDING MATER OFFSET PRINTING	RIAL A	D PRINTING PLATE PRODUCED THEREFROM FOR WATER	RLESS

#### (57) Abstract

The invention relates to a recording material having a plate—or sheet—like substrate, at least one IR—absorbing layer, which comprises at least one IR—absorbing component and at least one polymeric, organic binder and decomposes under the action of IR laser radiation or changes so that its adhesion to the silicone top layer decreases, and a top layer comprising a cured silicone rubber. The substrate comprises an oxidizable metal or one of its alloys and, at least on the side facing the IR—absorbing layer, is roughened and is covered with a layer of an oxide of the metal. The invention also relates to a process for the production of a waterless offset printing plate and to the printing plate itself which is produced from the recording material according to the invention and in which the hydrophilic surface of the oxide layer is ink—carrying and the oleophobic top layer is ink—repellent.

## FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
ΑU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of Americ
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Potand		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	LI	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

1

# LASER-IMAGE:ABLE RECORDING MATERIAL AND PRINTING PLATE PRODUCED THEREFROM FOR WATERLESS OFFSET PRINTING

#### DESCRIPTION

#### 1. Field of the invention.

The invention relates to a recording material having a plate- or sheet-like substrate, a top layer comprising a cured silicone rubber and at least one IR-absorbing layer, which comprises at least one IR-absorbing component and at least one polymeric, organic binder and decomposes under the action of IR laser radiation or changes so that its adhesion to the silicone top layer decreases. It can be provided with an image by means of laser radiation and serves primarily for the production of offset printing plates which print by a waterless method.

#### 2. Background of the invention.

Recording materials for waterless lithographic printing, where a laser-sensitive layer is removed, are already known. 12 038 describes a material which consists of a substrate, an intermediate layer which contains particles absorbing laser energy (in particular carbon black), nitrocellulose and a crosslinking agent, and a silicone rubber layer. Aluminium, paper and plastic are mentioned as substrates. To prevent the heat generated by the laser from being conducted away by the aluminium, the aluminium surface is preferably provided with an insulating layer of an oleophilic resin. The plate is exposed to infrared or visible laser radiation, preferably from an Nd-YAG laser or an argon laser. In the parts which the radiation strikes, the intermediate layer is oxidized and combusted. Consequently, the silicone layer present on top becomes detached and can be removed with an organic solvent. However, the recording material has only relatively little sensitivity. Moreover, the plates produced therefrom permit only a short print run.

EP-A 573 091 and EP-A 685 333 likewise describe a material for the production of waterless offset printing plates. Once again, it comprises a substrate having an oleophilic surface, a recording layer which is applied thereon, is not more than 3 μm thick and contains a substance which converts radiation into heat, and a cured

2

silicone layer. The substrates used are generally films of polyester, polycarbonate or polystyrene. Polyolefin-coated paper is also suitable. Aluminium substrates are also mentioned; however, these must be provided with a special oleophilic coating. Carbon black and pigments and dyes which absorb in the infrared range are mentioned as substances which convert radiation into heat. recording layer may also comprise a metal, e.g. bismuth, tin or tellurium, applied by vapour deposition. This metal layer is not more than 25 nm thick. The recording material is provided with an image by means of laser radiation and is then rubbed dry. During this procedure, the irradiated parts of the recording layer are removed together with the silicone layer present on top. Offset printing plates having a polyester substrate permit only relatively short runs in generally small-size, low-speed printing presses. potential uses of such printing plates are therefore greatly restricted.

EP-A 580 393 discloses, inter alia, a three-layer recording material for the production of waterless offset printing plates. comprises in general a substrate which reflects IR radiation, for example a substrate of degreased, bright-rolled aluminium or a polyester film, on which a reflecting aluminium layer has been applied by vacuum vapour deposition or by sputtering. absorbing layer and a silicone top layer are then applied to this substrate. The IR-absorbing layer is removed by imagewise exposure to laser radiation of appropriate wavelength. Consequently, those parts of the silicone layer which are present on top become detached and can be removed mechanically, for example by means of brushing. The metallic or metallized substrate has only little affinity to water. The disadvantage of such a recording material is once again the low adhesion between the substrate and the layer present on top. A printing plate produced therefrom accordingly gives only a short print run.

According to EP-A 644 047, a further layer which itself does not absorb laser radiation but, under the action of laser radiation on the IR-absorbing layer present on top, undergoes thermal decomposition with formation of gaseous products is arranged between the substrate and the IR-absorbing layer. The thickness of this further layer is chosen so that it undergoes only partial decomposition. In general, it is from 1 to 30  $\mu$ m thick. An adhesion-promoting layer, for example a layer of a silane or a protein, may also be arranged between substrate and thermally

3

decomposable layer.

#### 3. Summary of the invention.

It is therefore an object of the present invention to provide recording materials from which printing plates for waterless lithographic printing which have high sensitivity to infrared laser radiation and high resolution can be produced in a simple manner.

It is therefore a further object of the present invention to provide a printing plate which achieves a long print run.

According to the present invention there is provided a recording material having

- a plate- or sheet-like substrate,
- a top layer comprising a cured silicone rubber, and
- at least one IR-absorbing layer which comprises at least one IR-absorbing component and at least one polymeric, organic binder and decomposes under the action of IR laser radiation or changes so that its adhesion to the silicone top layer decreases, which is characterized in that the substrate comprises an oxidizable metal or one of its alloys and, at least on the side facing the IR-absorbing layer, is roughened and is covered with a layer of an oxide of the metal.

#### 4. Detailed description of the invention.

The substrate preferably consists of aluminium or one of its alloys. It is in general mechanically, chemically and/or electrochemically roughened. This roughening can be achieved by dry brushing, wet brushing, sand-blasting, chemical treatment and/or electrochemical treatment. The electrochemical roughening is preferred. It leads to outstanding anchoring of the IR-absorbing layer on top. The average peak-to-valley height  $\rm R_Z$  (determined according to DIN 4768 - October 1970 edition) of the surface is in the range from about 0.5 to 15  $\rm \mu m$ .

Suitable roughening methods are also described in EP-A 292 801, EP-A 437 761 and DE-A 33 05 067. The metallic substrate generally has high thermal conductivity. Its surface area increases owing to the roughening, with the result that the heat induced by the laser radiation can be removed even more rapidly. On the other hand, the metal oxide layer has a heat-insulating action and substantially

4

slows down the loss of heat. The layer comprising the metal oxide, especially alumina, has in general only 1/10 or less of the thermal conductivity of the respective metal. The oxide is preferably produced electrochemically directly from the metal of the substrate. Particularly in the case of aluminium substrates, the electrochemical oxidation can be controlled so that pores form in the oxide layer and even further reduce the thermal conductivity. The production of such oxide layers is generally known and described (for example in EP-A 161 461). As a result of the metal oxide layer, a hydrophilic, abrasion-resistant surface is produced on the substrate. Depending on the process parameters, the weight of the oxide layer is in general from 0.5 to 10  $g/m^2$ , preferably from 1 to  $5 \text{ g/m}^2$ . The surface obtained by the combination of roughening and oxidation reflects the IR laser radiation to a substantially lesser extent so that excellent reproduction of fine image elements and hence high resolution are ensured. The surface of the metal oxide layer is hydrophilic. Surprisingly, this hydrophilic surface shows excellent ink acceptance during subsequent printing. The prior art had suggested that only an oleophilic surface meets this requirement. The waterless offset printing plates produced from the recording material according to the invention give prints of excellent quality. The achievable print run is long; in general, it is more than 100,000 prints.

The IR-absorbing layer contains components, in particular pigments or dyes, which absorb laser radiation having a wavelength in the infrared range (especially in the range from 700 to 1200 nm). Here, the pigments are also to include carbon black. Suitable IR absorbers are mentioned in J. Fabian et al., Chem. Rev. 92 [1992] 1197. Pigments which contain metals, metal oxides, metal sulphides, metal carbides or similar metal compounds are also suitable. Finely divided metallic elements of main groups II to V and of subgroups I, II and IV to VIII of the Periodic Table, such as Mg, Al, Bi, Sn, In, Zn, Ti, Cr, Mo, W, Co, Ir, Ni, Pd, Pt, Cu, Ag, Au, Zr or Te, are preferred. Other suitable IR-absorbing components are metalphthalocyanine compounds, anthraquinones, polythiophenes, polyanilines, polyacetylenes, polyphenylenes, polyphenylene sulphides and polypyrroles. In order to avoid unnecessarily impairing the resolution, the absorbing pigment particles should have a mean diameter of, as far as possible, not more than 30  $\mu m$ . The amount of the IR-absorbing component is in general from 2 to 80% by weight, preferably from 5 to 57% by weight, based in each case on

5

the total weight of the nonvolatile components of the layer. IR-absorbing layer furthermore contains at least one polymeric, organic binder. Binders which undergo spontaneous decomposition under the action of heat are particularly advantageous. These binders undergoing autoxidation include in particular nitrocellulose. Polymers which do not undergo autoxidation and which undergo thermally induced decomposition indirectly with the formation of gaseous or volatile cleavage products may also be used. Examples of these are ethylcellulose, (meth)acrylate polymers and copolymers (such as poly(methyl methacrylate), poly(butyl acrylate), poly(2-hydroxyethyl methacrylate), copolymers of lauryl acrylate and methacrylic acid, polystyrene, poly(methylstyrene), copolymers of vinyl chloride and vinyl acetate, polyurethanes, polycarbonates and polysulphones. The directly or indirectly thermally decomposable polymers are not required in every case, so that other film-forming polymers may also be used. This applies when the IR-absorbing component already forms sufficiently volatile products under irradiation. For example, carbon black undergoes combustion when IR laser radiation strikes it, and accordingly gives gaseous combustion products. The "other film-forming polymers" are in particular homoand copolymers containing units of (meth)acrylic acid, (meth)acrylates and/or (meth)acrylamides, as well as polyvinyl acetates and polyvinyl acetals, which, if appropriate, are furthermore modified with carboxyl groups. They are used either in combination with the thermally decomposable materials or alone. The amount of the binders is in general from about 20 to 95% by weight, preferably from 30 to 80% by weight, based in each case on the total weight of the nonvolatile components of the layer.

In addition, the layer may also contain compounds which crosslink the binder. The type of crosslinking agent depends on the chemical functionality of the binder (S. Paul, Crosslinking Chemistry of Surface Coatings in Comprehensive Polymer Science, Volume 6, Chapter 6, page 149). The amount of the crosslinking agent or agents is in general from 0 to 30% by weight, preferably from 3 to 20% by weight, particularly preferably from 5 to 15% by weight, based in each case on the total weight of the nonvolatile components of the layer.

The IR-absorbing layer may moreover contain compounds which undergo decomposition under the action of heat and/or IR radiation or by chemical induction and form chemically active species (in particular acids), which in turn cause cleavage or decomposition of

6

the polymeric, organic binder. Once again, volatile cleavage or decomposition products are formed. Binders which contain tert-butoxycarbonyl groups give, for example, CO<sub>2</sub> and isobutene when acid acts thereon. Furthermore, the layer may contain compounds which form low molecular weight, gaseous or at least volatile cleavage products (Encycl. Polym. Sci. Eng., Vol. 2, page 434). Examples of such compounds are diazonium salts, azides, bicarbonates and azobicarbonates. The IR-absorbing layer can in addition contain stabilizers for increasing the storability, plasticizers, catalysts for initiating the crosslinking reaction, dulling agents, additional dyes, surfactants, levelling agents or other auxiliaries for improving stability, processing or reprographic quality. The amount of these additives is in general from 0 to 50% by weight, preferably from 10 to 30% by weight, based in each case on the weight of the nonvolatile components of the layer.

The weight of the IR-absorbing layer is in general from 0.15 to  $5~\rm g/m^2$ , preferably from 0.5 to  $3.5~\rm g/m^2$ . The IR-absorbing layer is produced by applying and drying an appropriate coating solution. Suitable solvents for the preparation of the coating solution include ketones, esters, glycol ethers, alcohols, ethers or mixtures thereof.

The silicone top layer is oleophobic and repels the ink during printing. It consists of crosslinked silicone rubber. Any silicone rubber which is sufficiently ink-repellent to permit printing without damping solution is in principle suitable. The term "silicone rubber" is intended to be understood as meaning a high molecular weight, essentially linear diorganopolysiloxane, in accordance with the definition of Noll "Chemie und Technologie der Silikone" [Chemistry and Technology of the Silicones], Verlag Chemie [1968], page 332. For the crosslinked or vulcanized products, on the other hand, the term "vulcanized silicone rubber" is used. In general, a solution of silicone rubber is applied to the IR-absorbing layer, dried and crosslinked. Particularly suitable solvents are toluene, xylene and in particular isoparaffins (boiling range from 100 to 180°C).

The silicone rubbers may be one-component or multicomponent rubbers. Examples are described in DE-A 23 50 211, 23 57 871 and 23 59 101. Particularly suitable one-component silicone rubbers are polydimethylsiloxanes which carry hydrogen atoms, acetyl, oxime, alkoxy or amino groups or other functional groups at the chain ends. The methyl groups in the chain may also be replaced by other alkyl

7

groups, by haloalkyl groups or by unsubstituted or substituted aryl groups (in particular phenyl groups). The terminal functional groups are readily hydrolysable and cure in a short time (from a few minutes to a few hours) in the presence of moisture. The multicomponent silicone rubbers are crosslinkable by addition or condensation. The addition-crosslinkable types contain in general two different polysiloxanes. One polysiloxane is generally present in an amount of from 70 to 99% by weight and has alkylene groups (in particular vinyl groups). The other is present in general in an amount of from 1 to 10% by weight. In this polysiloxane, hydrogen atoms are bonded directly to silicon atoms. The addition reaction is effected by heating to more than 50°C in the presence of from about 0.0005 to 0.002% by weight of a platinum catalyst. Multicomponent silicone rubbers have the advantage that they crosslink very rapidly at relatively high temperature (about 100°C). The time within which they can be processed, the so-called "pot life", is on the other hand often relatively short. The condensation-crosslinkable mixtures contain diorganopolysiloxanes having reactive terminal groups, such as hydroxyl and acetoxy groups. These are crosslinked with silanes or oligosilanes in the presence of catalysts. Crosslinking agents are present in an amount of from 2 to 15% by weight, while the catalysts are present in an amount of from 0.01 to 10% by weight, based in each case on the total weight of the nonvolatile components of the layer. These mixtures, too, react relatively rapidly and therefore have only a limited pot life.

A particularly preferred mixture consists of

- hydroxyl-terminated polydimethylsiloxanes,
- a silane crosslinking component (in particular a tetra- or trifunctional alkoxy-, acetoxy-, amido-, amino-, aminoxy-, ketoximino- or enoxysilane) or functionalized silicone resins,
- a crosslinking catalyst (in particular an organotin or organotitanium compound) and
- if appropriate, further components, such as
  - organopolysiloxane compounds having Si-H bonds,
  - platinum catalysts for further addition crosslinking,
  - silanes having adhesion-improving properties,
  - reaction inhibitors, '
  - fillers and/or dyes.

The stated silane crosslinking components and the reactions occurring during the crosslinking are described by J.J. Lebrun and

8

H. Porte in "Comprehensive Polymer Science", Vol. 5 [1989] 593-609.

The silicone layer may contain further components. These may serve for additional crosslinking, better adhesion, mechanical strengthening or colouring. These further components which are only optionally present may be present in an amount of not more than 10% by weight, preferably not more than 5% by weight, based in each case on the total weight of the layer.

After coating with the silicone rubber solution and drying, crosslinking ("curing") to give a vulcanized silicone rubber is carried out in a manner known per se by the action of moisture or spontaneously at room temperature or elevated temperature. The cured silicone rubber layer is virtually insoluble in organic solvents. It is transparent for the IR laser radiation and itself absorbs virtually no IR radiation (compounds with which the silicone layer may be coloured must therefore be appropriately chosen). The weight of the cured silicone layer is in general from 1 to 20  $g/m^2$ , preferably from 1 to 5  $g/m^2$ , which corresponds approximately to a thickness of from 0.8 to 17  $\mu$ m, preferably from 0.8 to 4  $\mu$ m.

The printing plates are produced by imagewise exposure to IR laser radiation. YAG lasers, Nd-YAG lasers, argon lasers, semiconductor lasers and laser diodes, each of which emit radiation in the IR range, are preferred. They generally have an output power of between 40 and 7500 mW. The radiation energy is in general from 20 to 600 mJ/cm<sup>2</sup>. As low a radiation energy as possible is desirable. The laser radiation results in ablation of the IR-absorbing layer, with the result that at the same time the oleophobic silicone layer becomes detached and is simultaneously removed. Loosely adhering layer components can be removed mechanically (for example by wiping), if appropriate with a suitable solvent. The printing plates can be produced in this manner in a single step. A further advantage is that only a very small amount of liquid waste products, if any, are obtained.

The present invention thus also relates to a process for the production of a waterless offset printing plate, which is characterized in that IR laser radiation is allowed to act imagewise on the recording material according to the invention. In the case of the waterless offset printing plate thus produced, the hydrophilic surface of the oxide layer is ink-carrying and the oleophobic top layer is ink-repellent.

The present invention finally also relates to the printing plate produced from the recording material according to the invention.

9

The examples which follow illustrate the invention. Therein, pbw represents part by weight. Unless stated otherwise, "%" represents "% by weight".

#### Example 1

A 0.3 mm thick, electrolytically roughened and anodically oxidized aluminium plate having an oxide weight of 3.6  $g/m^2$  was rendered hydrophilic with a 0.1% by weight aqueous polyvinylphosphonic acid solution. A solution of

44.4 pbw of a 20% by weight <sup>®</sup>Efweko NC 118 solution (Degussa AG), which consists of

18.0 pbw	of High-Color-Channel (HCC) carbon
	black,
28.0 pbw	of collodium wool of standard type 24
	Ε,
28.0 pbw	of collodium wool of standard type 27
	Ε,
22.0 pbw	of dibutyl phthalate and
4.0 pbw	of the copper salt of 2-ethylhexanoic
	acid in propylene glycol methyl ether
	acetate,

7.88 pbw of a 20% by weight solution of a mixture of diphenylmethane 4,4'-diisocyanate and polymeric components (\*\*Desmodur VKS) and

4.20 pbw of a 1% by weight solution of silicone oil in butan-2-one in a mixture of

73.2 pbw of butan-2-one and

20.3 pbw of propylene glycol methyl ether acetate (PGMEA)

was applied to this substrate and then dried for 2 min at 120°C. The weight of the resulting layer was  $3.0~{\rm g/m^2}$ .

#### A solution of

33.5 pbw of a 33% by weight solution of polydimethylsiloxanes containing hydroxyl groups in toluene (viscosity 9,000-15,000 mPa·s at 25°C, Wacker \*\*Dehesive 810),

1.56 pbw of a 50% by weight solution of silicone resin having aminoalkyl groups in toluene (Wacker V 83)

0.67 pbw of dibutyltin diacetate in

214 pbw of isoparaffin (initial boiling point 118°C)

10

was then applied to the IR absorber layer and thereafter dried for 2 min at 120°C with circulating air. The weight of the silicone layer was then  $2.3~{\rm g/m^2}$ .

The recording material thus produced was mounted on a uniformly driveable drum and recorded on by means of a continuously operating Nd-YAG laser (200 mW output power at 1064 nm) with a resolution of 1200 dpi. The exposure time, which was variable by changing the rotational speed of the drum, was set at 15 µs per pixel. This corresponds to an energy of about 500 mJ/cm<sup>2</sup>. The material on which various line patterns are recorded is wiped with a 1% strength by weight surfactant solution (e.g. <sup>®</sup>Glucopon 600 CS UP, Henkel) to remove the ablated material and is used for printing under the conditions usual for waterless offset. Well over 100,000 prints of excellent quality can be produced. The printing capacity is accordingly well above the usual prior art (in this context, cf. Seybold Report Vol. 24, No. 15-14 April, 1995, and Paper presented at New Era: A Technical Conference for the Printing Industry, held at Sutton Coldfield, UK, 16 March 1994, 17 pages).

#### Examples 2 to 10:

A 0.2 mm thick, electrolytically roughened and anodically oxidized aluminium plate having an oxide weight of  $3.4~\mathrm{g/m^2}$  was rendered hydrophilic with a 0.1% by weight aqueous polyvinylphosphonic acid solution. Coating solutions are prepared with the components stated in Table 1 and are applied, as described in Example 1, to this substrate type and dried.

Table 1: Components for layers absorbing IR radiation

PCT/EP98/00146

11

WO 98/31550

Component	2	3	4	5	6	7	8	9	10
Carbon									
black A <sup>1</sup> )	11.8	-	-	-	-	9.93	9.41	9.39	9.30
Carbon black B <sup>2)</sup>	_	7.05	_	_	_	_		_	_
Carbon	_	7.05							
black C3)	-	-	4.58	-	-	-	-	-	-
Carbon				7.25	15.0	_			
black D <sup>4)</sup> VP-N-3108 <sup>5)</sup>	]_	_	5.25	6.78	-	_	_	_	_
Carboset 526 <sup>6</sup> )	-	-	-	-	3.57	-	-	-	-
Cymel 301 <sup>7)</sup>									
(20% MEK)	3.13	1.88	2.63	2.63	2.63	-	-	-	2.63
Desmodur VKS <sup>8</sup> )									
(20% MEK)	-	-	-	-	-	-	5.25	-	-
Desmodur VK <sup>9</sup> )									
(20% MEK)	-	-	-	-	-	2.63	-	-	-
Silane Z- 6124 <sup>10</sup> )	-	-	-	-	-		-	_	0.53
EP 140 <sup>11)</sup> (20% MEK)	<b> </b> _	_	_	_	_		_	5.25	_
pTosOH <sup>12)</sup>								3.23	
(10% MEK) DABCO <sup>13)</sup>	1.25	0.79	1.05	1.05	1.05	-	-	-	1.05
(10% MEK)	-	-	-	-	-	_	-	0.21	-
Silicone	_		4.20	4.20		4.20	4.20	4.20	4.20
oil (1% MEK)	-	-	4.20	4.20	_	4.20	4.20	4.20	4.20
Ethyl								-	-
acetate	232	140	-	~	128	-	-		
Butyl	1.66	1.00	_	_	_	_	_		-
acetate 2-Butanone		-	76.5	76.5	_	77.4	75.3	75.2	76.5
Propylene				, , , ,			.5.5	,,,,	, , , ,
glycol methyl									
ether									
acetate			55.8	51.6		55.8	55.8	55.8	37.2
Layer									
weight [g/m <sup>2</sup> ]	1.6	2.2	4.1	3.9	3.0	2.8	2.2	2.6	2.4

- 1) Nitrocellulose chips (Degussa, Efweko NC 118/2)
- 2) Nitrocellulose pigment preparation (Hagedorn 71907)
- 3) Nitrocellulose pigment preparation (Hagedorn 70907)

12

6)	Polyacrylate resin (BF Goodrich)
7)	Melamine resin (Dyno Cyanamid)
8), 9)	Liquid polyisocyanates (Bayer)
10)	Phenyltrimethoxysilane (Dow Corning)
11)	Epoxy resin ( <sup>®</sup> Beckopox from Vianova Resins GmbH)
12)	p-Toluenesulphonic acid
13)	Diazabicyclooctane

The silicone rubber described in Example 1 is applied to the various IR absorber layers. The resulting printing plates are provided with an image analogously to Example 1, processed and used for printing.

### Examples 11 to 19:

An IR absorber layer is produced similarly to Example 1, covered with the silicone coating solutions shown in Table 2 and dried for 2 minutes at 120°C. The resulting layer weights are likewise listed in Table 2.

13

Table 2: Components for silicone layers

Component	11	12	13	14	15	16	17	18	19
Polymer1) Polymer2) Polymer3) Polymer4) Polymer5) V 836) Tetra(2-	18.8 - - - - 0.88	17.7 - - - - 0.88	17.7 - - - - 0.88	- 13.3 - - - 0.94	19.9 - - - -	- - 19.4 - -	- - - 9.46 -	- - - - 8.59	- - - - 8.41
methoxy- ethanol)- silane V 93 <sup>7)</sup> V 24 <sup>8)</sup> Ethyltri- acetoxysil	- - -	- - -	- - -	- - -	0.04	- 2.33 -	- - 0.24	0.32	-
Methyltris (methyleth yl- ketoximino )silane Dibutyltin diacetate OL <sup>9</sup> ) GF 9110) HF 8611) Iso- paraffin <sup>12</sup>	- 0.37 - - - 80.0	- 0.37 - 0.35 - 80.7	- [0.37 - - 0.35 80.7	0.40 - - - 85.3	- 0.20 - - - 79.8	- 0.06 - - 78.2	- 0.10 - 0.20	- 0.001 91.0	0.32 0.01 - 0.27 - 91.0
Layer weight [g/m <sup>2</sup> ]	4.3	4.1	4.3	2.8	3.8	2.9	2.1	2.5	2.3

- 1) 33% by weight solution of polydimethylsiloxanes containing hydroxyl groups in toluene (viscosity 9,000-15,000 mPa s at 25°C, Wacker Dehesive 810)
- 2) 50% by weight solution of polydimethylsiloxanes containing hydroxyl groups in toluene (viscosity 9,000-15,000 mPa·s at 25°C, Wacker Dehesive 850)
- 3) 30% by weight solution of polydimethylsiloxanes containing vinyl groups in naphtha, boiling range 80-110°C (viscosity 6,000-10,000 mPa·s at 25°C, Wacker Dehesive 940)
- 4) Polydimethylsiloxanes containing vinyl groups (viscosity

14

400-600 mPa·s, Wacker Dehesive 920)

- 5) Hydroxyl-terminated polydimethylsiloxanes (viscosity about 45,000 mPa·s, Bayer <sup>®</sup>Silopren E 50)
- 6) 50% by weight solution of a silicone resin having aminoalkyl groups in toluene (Wacker)
- 7) 5% by weight solution of a polyhydrogenomethylsiloxane in naphtha of boiling range 80-110°C (Wacker)
- 8) Polyhydrogenomethylsiloxane (viscosity 15-30 mPa·s at 25°C, Wacker)
- 9) 1% by weight solution of a platinum complex preparation in polydimethylsiloxane (Wacker)
- 10) [3-(2-Aminoethylamino)propyl]trimethoxysilane (Wacker)
- 11) Silane mixture comprising triacetoxyvinylsilane and trimethoxy(3-oxiranylmethoxypropyl)silane (Wacker)
- 12) Initial boiling point 118°C

The further processing of the printing plates is carried out as described in Example 1.

15

#### · CLAIMS

- 1. Recording material having
- a plate- or sheet-like substrate,
- a top layer comprising a cured silicone rubber, and
- at least one IR-absorbing layer which comprises at least one IR-absorbing component and at least one polymeric, organic binder and decomposes under the action of IR laser radiation or changes so that its adhesion to the silicone top layer decreases, and

which is characterized in that the substrate comprises an oxidizable metal or one of its alloys and, at least on the side facing the IR-absorbing layer, is roughened and is covered with a layer of an oxide of the metal.

- 2. Recording material according to Claim 1, characterized in that the substrate is mechanically, chemically and/or electrochemically roughened.
- 3. Recording material according to Claim 1 or 2, characterized in that the layer of the oxide is produced electrochemically.
- 4. Recording material according to Claim 3, characterized in that the oxide layer is porous.
- 5. Recording material according to any of Claims 1 to 4, characterized in that the weight of the oxide layer is from 0.5 to  $10~\mathrm{g/m^2}$ .
- 6. Recording material according to Claims 5, characterized in that the weight of the oxide layer is from 1 to 5  $g/m^2$
- 7. Recording material according to any of Claims 1 to 6, characterized in that the IR-absorbing component is a pigment, a dye, a metallic element or carbon black.
- 8. Recording material according to any of Claims 1 to 7, characterized in that the IR-absorbing layer contains a directly or indirectly thermally decomposable binder.
- 9. Recording material according to Claim 8, characterized in that SUBSTITUTE SHEET (RULE 26)

16

the directly thermally decomposable binder is nitrocellulose.

- 10. Recording material according to any of Claims 1 to 9, characterized in that the weight of the IR-absorbing layer is from 0.15 to 5  $g/m^2$ .
- 11. Recording material according to Claim 10, characterized in that the weight of the IR-absorbing layer is from 0.5 to 3.5  $\rm g/m^2$
- 12. Process for the production of a waterless offset printing plate, characterized in that IR laser radiation is allowed to act imagewise on the recording material according to any of Claims 1 to 11 and those parts of the IR-absorbing layer struck by the radiation are thus removed together with those parts of the top layer which are present on top.
- 13. Process according to Claim 12, characterized in that any adhering loose layer components are removed mechanically, optionally with the use of a solvent.
- 14. Waterless offset printing plate, characterized in that it is produced from a recording material according to any of Claims 1 to 11.

## INTERNATIONAL SEARCH REPORT

GO.

Inter inal Application No PCT/EP 98/00146

			01/21 30	, 001 10			
A. CLASSI IPC 6	FICATION OF SUBJECT MATTER B41M5/36 B41C1/10			·			
According to	o International Patent Classification (IPC) or to both national classific	ation and IPC					
	SEARCHED		•				
Minimum do IPC 6	ocumentation searched (classification system followed by classification B41M B41C	on symbols)	-				
Documentat	tion searched other than minimum documentation to the extent that s	uch documents are included	d in the fields sea	arched			
Electronic d	ata base consulted during the international search (name of data ba	se and, where practical, se	arch terms used)				
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT						
Category *	Citation of document, with indication, where appropriate, of the rele	evant passages		Relevant to claim No.			
Α	DE 25 12 038 A (SCOTT PAPER CO) 2 October 1975 cited in the application						
	see page 4, line 12 - line 21; clean example 2	laims;					
Α	PATENT ABSTRACTS OF JAPAN vol. 015, no. 214 (P-1209), 31 May 1991 & JP 03 059558 A (KONICA CORP; OTHERS: 01), 14 March 1991, see abstract						
A	WO 94 01280 A (DU PONT UK ;GATES PETER (GB); FELDER THOMAS CLARK ( January 1994 see example 3						
:		,					
	-	-/					
				İ			
X Furth	er documents are listed in the continuation of box C.	X Patent family men	nbers are listed in	n annex.			
* Special cat	egories of cited documents :	"T" later document publish					
conside	nt defining the general state of the art which is not ared to be of particular relevance ocument but published on or after the international	or priority date and no cited to understand th invention	e principle or the	ory underlying the			
filing da		"X" document of particular cannot be considered involve an inventive s	novel or cannot	aimed invention be considered to sument is taken alone			
which t citation	s cited to establish the publication date of another or other special reason (as specified)	"Y" document of particular cannot be considered	relevance; the cl	almed invention entive step when the			
other n		document is combined ments, such combination in the art.	d with one or mo	re other such docu-			
"P" docume later th	nt published prior to the international filing date but an the priority date claimed	*& document member of t	he same patent f	amily			
Date of the s	ctual completion of theinternational search	Date of mailing of the i	nternational sear	ch report			
27	7 May 1998	08/06/199	8				
Name and m	nailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2	Authorized officer					
	NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl,	Philosoph	ı. <b>1</b>				

1

## INTERNATIONAL SEARCH REPORT

Interr nal Application No PCT/EP 98/00146

		PCT/EP 98/00146	
	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	72.	
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to clain	n No.
A	DATABASE WPI Section Ch, Week 9133 Derwent Publications Ltd., London, GB; Class A26, AN 91-243502 XP002066046 & JP 03 159 793 A (TOPPAN PRINTING CO LTD) see abstract		
A	EP 0 580 393 A (PRESSTEK INC) 26 January 1994 cited in the application see page 9, line 27 - line 41; example 13A see page 15, line 16 - line 41; figures 13A,13I; examples 8-10,18		
P,A	DATABASE WPI Section Ch, Week 9808 Derwent Publications Ltd., London, GB; Class A89, AN 98-081721 XP002066047 & JP 09 315 024 A (FUJI PHOTO FILM CO LTD) see abstract		

1

INTERNATIONAL SEARCH REPORT

autormation on patent family members

Intern. .nal Application No PCT/EP 98/00146

Patent docume cited in search re		Publication date	i	Patent family member(s)	Publication date
DE 2512038	A .	02-10-1975	CA	1050805 A	20-03-1979
			FR	2264671 A	17-10-1975
			GB	1489308 A	19-10-1977
			JP	50158405 A	22-12-1975
WO 9401280	) A	20-01-1994	DE	69301826 D	18-04-1996
			DE	69301826 T	12-09-1996
			ĒΡ	0649374 A	26-04-1995
			ES	2084505 T	01-05-1996
			GB	2283703 A,B	17-05-1995
			JP	7508474 T	21-09-1995
EP 0580393	3 A	26-01-1994	US	5339737 A	23-08-1994
			AU	688702 B	12-03-1998
			AU	1017397 A	15-05-1997
			AU	1240697 A	15-05-1997
			AU	4178393 A	27-01-1994
			CA	2100517 A	21-01-1994
			JP	6199064 A	19-07-1994
			US	5353705 A	11-10-1994
			US	5379698 A	10-01-1995
			US	5487338 A	30-01-1996
			US	RE35512 E	20-05-1997
			AU	4178493 A	27-01-1994
			AU	6447996 A	07-11-1996
			CA	2100413 A	21-01-1994
			· EP	0580394 A	26-01-1994
			JP	2648081 B	27-08-1997
			JP	6186750 A	08-07-1994
·			บร	5351617 A	04-10-1994
			US	5385092 A	31-01-1995
			US	5540150 A	30-07-1996
			US	5551341 A	03-09-1996
			US	5638753 A	17-06-1997